

CORRECTION

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Correction: Curvature-induced hydrophobicity at imogolite–water interfaces

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Correction for ‘Curvature-induced hydrophobicity at imogolite–water interfaces’ by Alejandro Fernandez-Martinez et al., *Environ. Sci.: Nano*, 2020, DOI: 10.1039/d0en00304b.

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The corrected Acknowledgements section is shown below:

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The Royal Society of Chemistry apologises for these errors and any consequent inconvenience to authors and readers.

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